NTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 6: C12N 15/82, 15/12, 15/62, 5/10, C07K

(11) International Publication Number: A1

WO 98/21348

14/485, A01H 15/00

(43) International Publication Date:

22 May 1998 (22.05.98)

(21) International Application Number:

PCT/US97/20603

(22) International Filing Date:

12 November 1997 (12.11.97)

(30) Priority Data:

08/747.246

12 November 1996 (12.11.96) US

(71) Applicant: BATTELLE MEMORIAL INSTITUTE [US/US]; Pacific Northwest Division, Intellectual Property Services, P.O. Box 999, Richland, WA 99352 (US).

(72) Inventors: HOOKER, Brian, S.; 2525 W. Grand Ronde #14, Kennewick, WA 99336 (US). DAI, Ziyu; Apartment D-5. 1621 George Washington Way, Richland, WA 99352 (US). GAO, Jianwei; 506 Snyder Road, Richland, WA 99352 (US). KINGSLEY, Mark, T.; 531 North Reed Street, Kennewick, WA 99336 (US). WELLER, Richard, E.: 50 Herlou Place, Selah, WA 98942 (US).

(74) Agent: ZIMMERMAN, Paul, W.; Battelle Memorial Institute, Pacific Northwest Division, Intellectual Property Services, (K1-53), P.O. Box 999, Richland, WA 99352 (US).

(81) Designated States: CA, JP, European patent (AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE).

Published

With international search report.

Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

(54) Title: METHOD OF PRODUCING HUMAN GROWTH FACTORS FROM WHOLE PLANTS OR PLANT CELL CULTURES

(57) Abstract

The production of hEGF is achieved in both whole plants and plant cell culture wherein the hEGF has a length of at least 200 amino acids. For epidermal growth factor this would comprise at least a tetramer of EGF units. Effectiveness or production of the translation process has been increased according to the present invention by (1) cloning of pre-pro-EGF cDNA of approximately 4.5 kb into both whole plants and cell culture to increase overall titers of active hEGF; (2) synthesizing cDNA and transforming plants and cell culture for production of an oligomeric polypeptide consisting of repeated hEGF domains; and (3) increasing the overall size of the gene to be expressed with a fusion construct encoding hEGF linked to a protein that is efficiently produced in plant systems. As needed, synthetic cDNA includes plant-specific proteolytic cleavage sites between EGF repeats to facilitate correct processing in planta. Appropriate proteolytic cleavage sites upstream and downstream of hEGF are added if needed to obtain final product. In whole plants, use of a regulatory element confers hEGF production characteristics into traditionally non-saleable portions of crop plants, such as the leafy tops of potatoes. Use of potato tops under post-harvest conditions, results in overexpression production of hEGF in non-saleable plant portions towards the end of the harvesting season, without affecting crop quality.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL	Albania	ES	Spain	LS	Lesotho	SI	Slovenia
AM	Armenía	FI	Finland	LT	Lithuania	SK	Slovakia
AT	Austria	FR	France	LU	Luxembourg	SN	Senegal
ΑU	Australia	GA	Gabon	LV	Latvia	SZ	Swaziland
AZ	Azerbaijan	GB	United Kingdom	MC	Monaco	ΦT	Chad
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
BB	Barbados	GH	Ghana	MG	Madagascar	TJ	Tajikistan
BE	Belgium	GN	Guinea	MK	The former Yugoslav	TM	Turkmenistan
BF	Burkina Faso	GR	Greece		Republic of Macedonia	TR	Turkey
BG	Bulgaria	HU	Hungary	· ML	Mali	TT	Trinidad and Tobago
BJ	Benin	1E	Ireland	MN	Mongolia	UA	Ukraine
BR	Brazil	IL	Israel	MR	Mauritania	UG	Uganda
BY	Belarus	IS	Iceland	MW	Malawi	US	United States of America
CA	Canada	IT	Italy	MX	Mexico	UZ	Uzbekistan
CF	Central African Republic	JP	Japan	NE	Niger	VN	Viet Nam
CG	Congo	KE	Kenya	NL	Netherlands	YU	Yugoslavia
CH	Switzerland	KG	Kyrgyzstan	NO	Norway	zw	Zimbabwe
CI	Côte d'Ivoire	KP	Democratic People's	NZ	New Zealand		
CM	Cameroon		Republic of Korea	PL	Poland		
CN	China	KR	Republic of Korea	PT	Portugal		
CU	Cuba	KZ	Kazakstan	RO	Romania		
CZ	Czech Republic	LC	Saint Lucia	RU	Russian Federation		
DE	Germany	LI	Liechtenstein	SD	Sudan		
DK	Denmark	LK	Sri Lanka	SE	Sweden		
EE	Estonia	LR	Liberia	SG	Singapore		

WO 98/21348 PCT/US97/20603

METHOD OF PRODUCING HUMAN GROWTH FACTORS FROM WHOLE PLANTS OR PLANT CELL CULTURES

FIELD OF THE INVENTION

The present invention relates generally to a method for producing human growth factors from whole plants or plant cell culture. More specifically, the invention relates to producing a human growth factor from a plant cell encoded to produce the human growth factor with a length of at least 200 amino acids from transgenic plant cells.

BACKGROUND OF THE INVENTION

Growth factors and monoclonal antibodies (Mabs) are diverse yet highly specialized types of proteins having research and commercial applications in areas of therapeutics and diagnostics.

Therapeutic uses of human epidermal growth factor (hEGF) include treatment of soft tissue wounds (U.S. 5,218,093, 1993), specifically including skin and eye injuries as well as corneal and stomach ulcers (Frost and Sullivan 1996, 1994). In addition, several hEGF-bearing fusion constructs have been considered and/or tested, including mitotoxins for treatment of restenosis (Frost and Sullivan, 1994) and radioconjugates for a variety of anti-neoplastic therapies (Grieg et al., 1988).

Current production techniques for these proteins such as hybridoma and other types of mammalian cell culture methods (Köhler and Milsten, 1975) are generally slow, labor intensive, and consequently, expensive. In addition, current production techniques are difficult to validate due to the pathogenic and oncogenic potential of cultivated mammalian tissue.

Multimers of from 2 to 7 EGF units each having 53 amino acid residues have been produced from bacterial hosts, eg E. coli, Streptomyces and Bacillus, fungal hosts, eg Saccharomyces, Pichia and Aspergillus, insect cell host, and

5

10

15

20

25

10

15

20

25

mammalian cell hosts, eg CHO cells and COS cells. (U.S. Patent No. 5218093, 1993). hEGF production in *Staphylococcus aureus* (U.S. Patent No. 5004686, 1991) is by a fusion construct encoding hEGF linked to a protein. Synthesis methods using transgenic bacterial strains have problems such as faulty antibody gene expression, protein folding difficulties, inability to glycosylate proteins, and relegation of foreign peptides to insoluble material accumulated in inclusion bodies.

Transgenic plants can be used for the production of high value, medicinally important proteins, for example, production of Mabs (Hiatt et al., 1989; Düring et al., 1990; Benvenuto et al. 1991, Firek et al. 1993, Gao et al. 1993), human growth hormone (Kay et al. 1986) and human serum albumin (Sijmons et al. 1990). Transformed cells synthesize, secrete, and accumulate functional antibodies including single (Benvenuto et al. 1991) and double (Düring et al. 1990, Hiatt et al. 1991) domain immunoglobulins. However, it is noted that none of these authors investigated production of any human growth factor from transgenic plants.

Plant cell culture media are well-defined and inexpensive compared to mammalian cell culture media. Further, plant cell products, unlike mammalian-derived protein formulations, are generally assumed as neither pathogenic nor oncogenic to humans (Crawford, 1995). Also, when compared to similar production in transgenic bacterial strains (Attaai and Shuler 1987), plant tissue culture methods showed greater stability of foreign gene expression, even without use of selection pressure (Gao et al. 1991). One author, Higo et al. (1993) produced a human growth factor, specifically hEGF in transgenic tobacco with cDNA fragment size of 180 bp. Unsatisfactory foreign peptide levels of 20 to 60 pg/mg (ppb) total soluble leaf protein were obtained. This is despite the fact that plant progeny appeared to produce high levels of hEGF mRNA. Exact reasons for low observed levels of hEGF production are unclear. However, no signal peptide was encoded upstream of hEGF cDNA which could cause the foreign protein to be

10

15

relegated to the cytosol. Within this cell fraction, hEGF suffers proteolytic attack, especially considering the relatively small size (53 amino acids) or the peptide.

Although advantages have been observed for deriving proteins including EGF from plants, no transgenic plant cell culture process has been commercially developed for production of human growth factor. The lack of commercial exploitation of plant derived proteins is due in part to existing technological hurdles as observed by Higo et al. In addition, Ma et al., 1995 reported Mab titers of up to 500 µg/g (ppm) fresh weight of plant material (or 300 mg/L on a cell culture basis) whereas comparable mammalian cell processes are reported to attain levels of 1-2 g/L and higher (Rosenberg, personal communication, 1995). Implementation of alternative production systems to mammalian and bacterial culture, such as plant cellular techniques, has been further limited by non-technological factors, such as industry and regulatory acceptance (Simonsen and McGrogan, 1994) because of the investment made in developing and validating the more established non-plant methods.

Accordingly, there is a continuing need for plant based production of human growth factors.

Background References

20

- 1. U.S. Patent No. 5218093, 1993.
- 2. Frost & Sullivan. Emerging wound management technologies: wound healing/wound closure growth. February 1996.

- 3. Frost & Sullivan. World prescription dermatology pharmaceuticals markets. June 1994.
- 4. Grieg R, Dunnington D, Murthy U, Anzano M. 1988. Growth factors as novel therapeutic targets in neoplastic disease. Cancer Surveys 7(4):653-674.
 - 5. Köhler G, Milsten C. 1975. Continuous cultures of fused cells secreting antibody of predefined specificity. *Nature*. 256:495.
- 35 6. U.S. Patent No. 5004686, 1991.

- 7. Hiatt A, Cafferkey R, Bowdish K. 1989. Production of antibodies of transgenic plants. *Nature*. 342:76-78.
- 8. Düring K, Hippe S, Kreuzaler F, Shell J. 1990. Synthesis and self-assembly of a functional monoclonal antibody in transgenic *Nicotiana tabacum*. *J Plant Mol Biol* 15:281-293.
- 9. Benvenuto E, Ordas RJ, Tavazza R, Ancora G, Biocca S, Cattaneo A, Galeffi P. 1991. 'Phytoantibodies': a general vector for the expression of immunoglobulin domains in transgenic plants. *Plant Mol Biol* 17:865-874.
 - 10. Firek S, Draper J, Owen MRL, Gandecha A, Cockburn B, Whitelam GC. 1993. Secretion of a functional single-chain Fv protein in transgenic tobacco plants and cell suspension cultures. *Plant Mol Biol* 23:861-870.
 - 11. Gao J, Konzek RL, Linzmeier M, Buckley KB, Magnuson NS, Reeves R, An G, Lee JM. Production of monoclonal antibodies in plant cell culture. Presented at the 1993 ACS Fall Symposium, Denver.
- 20 12. Kay R, Chan A, Dayly M, McPherson J. 1987. Duplication of CaMV 35S promoter sequences creates a strong enhancer for plant genes. Science 236:1299-1302.
- Sijmons PC, Dekker BMM, Schrammeieijer B, Verwoerd TC, van den Elzen
 JM, Hoekema A. 1990. Production of correctly processed human serum albumin in transgenic plants. *Bio/Technology* 8:217-221.
 - 14. Crawford M. 1995. Therapeutic protein production using plant cell culture. Report by Lasure and Crawford, Inc.
 - 15. Attaai MM, Shuler ML. 1987. A mathematical model for predicting of plasmid copy number and genetic stability in Escherichia coli. *Biotechnol Bioeng* 30:389-397.
- 35 16. Gao J, Lee JM, An G. 1991. The stability of foreign protein production in genetically modified plant cells. *Plant Cell Reports* 10:533-536.
- Higo K, Saito Y, Higo H. 1993. Expression of a chemically synthesized gene for human epidermal growth factor under the control of cauliflower mosaic virus 35S promoter in transgenic tobacco. *Biosci Biotechnol Biochem* 57:1477-1481.

- Ma JKC, Hiatt A, Hein M, Vine N, Wang F, Stabila P, van Dolleweerd C, Mostov K, Lehner T. 1995. Generation and Assembly of Secretory Antibodies in Plants. Science 268:716-719.
- 5 19. Simonsen CC, McGrogan M. 1994. The molecular biology of production cell lines. *Biologicals* 22:85-94.

SUMMARY OF THE INVENTION

Despite the hurdles in technology development and commercialization, economic analysis indicates that regulatory costs associated with plant cell culture may reduce by as much as \$70,000 per batch as compared to analogous mammalian cell processes (Crawford, 1995). In addition, direct production costs for whole plant processes at equal protein production rates appear to be two to four orders-of-magnitude lower than comparable mammalian cell processes (Agracetus 1995). Additionally, as plant cell titers increase, this type of production becomes even more capital cost-effective.

It is, therefore, an object of the present invention to provide whole plant and plant cell culture derived human growth factors at higher overall concentrations and production rates, comparable to mammalian host cell systems.

It is a further object of the present invention to synthesize specific human growth factors.

It is another object of the present invention to increase production rates and concentrations by increasing protein stability through the use of fusion constructs.

It is a further object of the present invention to use *Nicotiana tabacum* (tobacco) and *Solanum tuberosum* (potato) whole plants and highly synchronous suspensions.

According to the present invention, the production of human growth factors is achieved in whole plants or plant cell culture wherein the human growth factor is produced with a length of at least 200 amino acids. For epidermal growth factor this would comprise at least a tetramer of EGF units.

20

25

10

15

20

Modifying chimeric cDNA and subcloning into a plant expression vector are done using standard molecular cloning procedures (Ausubel et al. 1992) and splicing PCR techniques (Marks et al. 1992).

Effectiveness or production of the translation process has been increased according to the present invention by (1) cloning of pre-pro-EGF cDNA of approximately 4.5 kb into both whole plants and cell culture to increase overall titers of active hEGF, (2) synthesizing cDNA and transforming plants and cell culture for production of an oligomeric polypeptide consisting of repeated hEGF domains, and (3) increasing the overall size of the gene to be expressed with a fusion construct encoding hEGF linked to a protein that is efficiently produced in plant systems. As needed, synthetic cDNA includes plant-specific proteolytic cleavage sites between EGF repeats to facilitate correct processing *in planta*. Appropriate proteolytic cleavage sites upstream and downstream of hEGF are added if needed to obtain final product.

The subject matter of the present invention is particularly pointed out and distinctly claimed in the concluding portion of this specification. However, both the organization and method of operation, together with further advantages and objects thereof, may best be understood by reference to the following description taken in connection with accompanying drawings wherein like reference characters refer to like elements.

BRIEF DESCRIPTION OF THE DRAWINGS

- FIG. 1 provides the size of EGF precursor (pre-pro-EGF) relative to correctly processed EGF.
 - FIG. 2 depicts schematically the construction of pZD203, a vector used to modify the restriction sites on pre-pro-EGF to develop cDNA suitable for cloning into the plant expression vector pGA643.
- FIG. 3 depicts schematically the construction of pZD204, the plant expression vector carrying pre-pro-EGF.

10

15

20

25

FIG. 4 shows EGF levels seen in individual calli resulting from positive transformation and antibiotic selection. EGF concentrations were determined using enzyme-linked immunosorbent assay and are based on a 30 KD protein size.

DESCRIPTION OF THE PREFERRED EMBODIMENT(s)

The present invention is a method for production of human growth factors using whole plants as well as plant cell suspensions transformed with appropriately constructed vector plasmids, wherein the human growth factor is produced with a length of at least 200 amino acids. More specifically, the method of the present invention is stable expression of human growth factors of interest as direct therapeutics, targeted delivery systems and research reagents. Human growth factors produced include human epidermal growth factor (hEGF), transforming growth factor (TGF), vascular endothelial growth factor (VEGF), platelet-derived growth factor (PDGF), fibroblast growth factor (FGF), tumor necrosis factor (TNF), heparin-binding epidermal growth factor (HBEGF), insulin-like growth factor (ILGF), platelet-derived endothelial cell growth factor (PDECGF), platelet-derived angiogenesis factor (PDAF), and bone-and-cartilage inducing growth factor (BCIF).

Any plant from the plant kingdom may be utilized. Specific types of plants that are amenable to the transformation steps listed herein include, but are not limited to monocotyledonous, dicotyledonous, and tuberous plants. Preferred species include but are not limited to *Nicotiana tabacum* (tobacco), *Solanum tuberosum* (potato), *Glycine max* (soybean), and *Zea mays* (corn).

The method of the present invention, a method of producing human growth factors from plant cells, has the steps of:

- (a) obtaining a positive transformant of the plant cells, the positive transformant carrying genetic material encoding the production of a human growth factor with a length of at least 200 amino acids;
- 30 (b) cultivating the positive transformant; and

10

15

20

(c) obtaining the human growth factors.

The step of obtaining may be as simple as purchasing or more complex actual making by well known methods, for example direct particle bombardment as described in Gene Transfer by Particle Bombardment, Klein TM, Knowlton S, Arentzen R, Plant Tissue Culture Manual, D1, pp 1-12, 1991, Kluwer Academic Publishers, or by Agrobaterium mediated transformation as described in Hoekema et al. 1985 (Hoekema KM, Hirsch PR, Hooykaaf PJJ, Schliperoort RA, 1985, Nononcogenic Plant Vectors for Use in the Agrobacterium Binary System, Plant Molecular Biology, Vol. 5, 85-89), and further described herein.

The step of cultivating involves either whole plant cultivating or tissue cultivating by any of well known cultivating methods.

The step of obtaining is by well known separation purification steps, for example ultrafiltration, affinity chromatography, and/or electrophoresis.

An Agrobacterium mediated transformation method of the present invention has the steps of:

- (a) modifying chimeric cDNA encoding a specific growth factor for subcloning into a plant expression vector
 - (b) subcloning the chimeric cDNA into the plant expression vector;
- (c) transferring the plant expression vector containing transgenic plant cells to an agrobacterium;
- (d) co-cultivating a portion of the transgenic plant cells (suspension culture or leaf disks) with the agrobacterium;
- (e) selecting positive transformants from the co-cultivated culture on an antibiotic selective media;
- 25 (f) permitting growth of the transgenic plant cells in whole plants or suspensions; and
 - (g) extracting a liquid containing the human growth factor; wherein the improvement comprises:

said human growth factor having a length of at least 200 amino acids.

10

15

20

25

Modifying chimeric cDNA and subcloning into a plant expression vector are done using standard molecular cloning procedures (Ausubel et al. 1992) and splicing PCR techniques (Marks et al. 1992). More specifically, modifying chimeric cDNA, has the steps of:

- (a) adding a transcription promoter to the upstream or 5' end of the chimeric cDNA; and
- (b) adding a transcription terminator to the downstream or 3' end of the chimeric cDNA. The transcription promoter and the transcription terminator are regulatory elements.
- Further, an additional regulatory element encoding a signal peptide may be added between the transcription promoter and the 5' end of the chimeric cDNA in order to relegate the product human growth factor to a specific cellular organelle. In addition, other regulatory elements may be added either between the promoter and the additional regulatory element encoding the signal peptide or at the 3' end of the chimeric cDNA to obtain greater mRNA stability between transcription and translation events.

In either whole plants or cell cultures, to enhance expression of the chimeric gene (hEGF), the present invention further includes manipulation of a 35S promoter by duplication of the upstream region (-343 to -90 bp) of the CaMV 35S promoter to increase transcription activity, as well as use of TSC29 and TSC40 promoters. These promoters and their transcription activity have been reported by Gao et al. 1994, and Dai et al. 1995.

In whole plants, transcription promoters may include the upstream enhancer (nucleotides -343 to -90 relative to the transcription start site) of the CaMV 35S promoter (Benfey et al. 1989) or the chlorophyll a/b binding protein (cab1) promoter (Ha and An 1988). Use of these types of regulatory elements confers human growth factor production characteristics into traditionally non-salable portions of crop plants, such as the leafy tops of potatoes. Use of potato tops, for example, under post-harvest conditions, results in overexpression and production

10

15

20

25

30

of human growth factor in non-salable plant portions towards the end of the harvesting season, without affecting crop quality.

Transferring the plant expression vector into the agrobacterium is completed using the freeze-thaw method (An 1987). For monocotyledonous species, superbinary vectors, such as pTOK233 and pSB131, are used to achieve high transformation frequency (Ishida et al. 1996). Remaining cocultivation, selection, growth, and extraction steps (d through g) have been described by Magnusen et al. (1996), and are well known in the art of plant molecular biology.

Many human growth factors possess relatively short lengths of between 50 and 100 amino acids. For example, hEGF has a length of 53 amino acids. Accordingly, obtaining a larger construct of at least 200 amino acids requires either (1) cloning the larger precursor cDNA, (2) synthesizing a concatemer consisting of multiple gene copies encoding the growth factor, or (3) increasing the overall size of a gene to be expressed using a fusion construct encoding a growth factor linked to a protein that is efficiently produced in plant systems.

An example of obtaining a larger precursor to increase the overall protein size is the cDNA encoding pre-pro-EGF. This particular gene, at approximately 4.5 kb, encodes a 1207 amino acid protein that, *in vivo*, is proteolytically cleaved to yield 53 amino acid EGF. In plant systems, this larger protein will provide additional stability against proteolytic degradation.

Synthesizing the cDNA concatemer is preferably done by ligating multiple gene copies using peptide linkers to obtain a processed protein length of at least 200 amino acids. The multiple gene copies are preferably an oligomeric polypeptide having of repeated growth factor cDNA domains. Peptide linkers may be used that are (1) proteolytically cleaved *in planta*, (2) proteolytically cleaved in a separate enzymatic treatment step, or (3) resistant to proteolytic cleavage. Peptide linkers that are proteolytically cleaved by serine proteases *in planta* preferably possess the amino acid sequence Arg-Asn. This sequence already exists when EGF is concatemerized since the C-terminal amino acid is arginine and the N-terminal amino acid is asparagine. To achieve *in planta* cleavage, the processed

10

15

20

25

30

protein is targeted either to the cell cytosol (no signal peptide) or vacuole (phytohemagglutinin signal peptide [Chrispeels et al.1991]). To achieve proteolytic cleavage in a separate enzymatic treatment step, the same amino acid sequence is preferably used (Arg-Asn) and the growth factor concatemer is either targeted to the chloroplast (pea photosystem II signal peptide) or secreted (PR-II signal peptide) to limit proteolytic degradation. To achieve resistance to proteolytic cleavage, linkers would preferably possess the amino acid sequence Arg-Pro. This sequence is resistant to serine proteases. Specifically for EGF, linkage would preferably be achieved by synthesizing cDNA encoding a single proline unit between growth factor monomers cDNA.

Increasing the overall size of a gene may be done by ligating EGF with cDNA encoding a protective protein to protect from proteolytic cleavage, thereby forming a fusion construct. Protective proteins include but are not limited to streptococcal protein G or -galactosidase, that have both been shown to inhibit proteolysis when attached to the C-terminus of other foreign proteins (Hellebust et al. 1989). Gene size could also be increased by ligating EGF with cDNA encoding another protective protein of commercial interest that processes well in plant-based systems. Protective proteins further include human serum albumin (Sijmons et al. 1990) and phytase (Verwoerd et al. 1995).

At least one genetic regulatory element may be included in the cDNA encoding the transcription of specific growth factors. Regulatory elements include transcription promoters or enhancers that increase the frequency of transcription events, leader sequences that increase the stability of mRNA prior to translation, and signal peptides that target proteins to specific organelles for posttranslational modifications and accumulation. Examples of transcription enhancers include but are not limited to the octapine synthase enhancer, a 16 bp palindrome (ACGTAAGCGCTTACGT) (Ellis et al. 1987) and the B-domain of the cauliflower mosaic virus 35S promoter (Kay et al. 1987). An example of a leader sequence includes but is not limited to alfalfa mosaic virus RNA4 leader sequence (Jobling and Gehrke 1987). Examples of signal peptides include but are not

limited to the tobacco PR-S signal peptide (Cornelissen et al. 1986) and the phytohemagglutinin signal peptide (Hunt and Chrispeels 1991).

Example 1

5

10

15

20

25

30

The bacteriophage λEGF116 (ATCC No. 59956) containing the gene encoding the full length polypeptide of human kidney pre-pro-EGF was obtained from ATCC. Pro-EGF (FIG. 1) is the 1207 amino acid precursor in which hEGF is flanked by polypeptide segments of 907 and 184 residues at its NH₂- and COOH-termini, respectively (Bell et al., 1986). The remainder of the 4.8 kb pre-pro-EGF gene encodes native signal peptides at both the NH₂- and COOH- termini of pro-EGF. The polypeptide contains a transmembrane (TM) binding region that facilitates proper cleavage in the endoplasmic reticulum.

The full length of cDNA was excised with Sma I, Hind III, and Eco RI restriction enzymes, as shown on FIG. 2, producing two separate fragments. These were sequentially ligated into compatible Sma I and Eco RI sites in pBluescript- creating the 7.5 kb plasmid pZD203. After proper orientation was confirmed, pre-pro-EGF cDNA was further excised with Xba I and Cla I restriction enzymes and ligated into compatible sites located between the CaMV 35S promoter and T₂ transcription terminator of binary vector pGA643, forming the 16 kb plasmid pZD204 (FIG. 3). This plasmid was directly transferred into Agrobacterium tumefaciens LBA4404 using the freeze-thaw method (An 1987). The transferred plasmid was introduced into tobacco whole plants (by leaf disks) and calli (by suspension culture) by co-cultivation with the Agrobacterium thereby producing transformants. Over 200 specific samples of transformants were taken from the co-cultivation and separately placed on kanamycin selective media. The co-cultivated transformants that grew were positive transformants. The positive transformants were screened under kanamycin selection pressure and preliminary ELISA results indicated the presence of hEGF in tobacco calli. Accumulation levels of hEGF in select transgenic calli are shown on a ng/g fresh weight basis in FIG. 4. The bars in FIG. 4 represent a random sample of the specific samples of transformants. The highest level of accumulation at approximately 400 ng/(g fresh weight cells) (ppb) corresponds to a concentration of 4.1 ng/(mg total soluble protein) (ppm) (based on a measured total soluble protein level of approximately 98 mg/(g fresh weight cells)). The 4.1 ng/(mg total soluble protein) (ppm) corresponds to 4100 pg/(mg total soluble protein) (ppb) which is almost two orders-of-magnitude greater than the result of 60 pg/(mg total soluble protein) (ppb) reported by Higo et al. (1993).

Further ELISA and Northern blot analyses were used to detect high levels of foreign protein production and mRNA transcription, respectively. Western blot analysis, completed to determine protein size, showed that specific EGF bearing constructs of 30 KD were produced. This size corresponds to approximately 250 amino acids.

Closure

While a preferred embodiment of the present invention has been shown and described, it will be apparent to those skilled in the art that many changes and modifications may be made without departing from the invention in its broader aspects. The appended claims are therefore intended to cover all such changes and modifications as fall within the true spirit and scope of the invention.

20

10

15

References

- 1. Crawford M. 1995. Therapeutic protein production using plant cell culture. Report by Lasure and Crawford, Inc., Seattle, Washington.
- Tuan J. 1995. Plant bioreactor systems program overview. Report by Agracetus, Inc., Middleton, Wisconsin.
- 3. Benfey PN, Ren L, Chua NH. 1989. The CaMV 35S enhancer contains at least two domains which can confer different developmental and tissue-specific expression patterns. *EMBO J* 8:2195-2202.
- Ha SB, An G. 1988. Identification of upstream regulator elements in the developmental expression of the Arabidopsis thaliana cab1 gene. *Proc Natl Acad Sci USA* 85:8017-8021.

- 14 -

- 5. Gao J, Kim SR, Chung YY, Lee JM, An G. 1994. Developmental and environmental regulation of ribosomal protein genes in tobacco. Plant Mol Biol 25:761-770.
- 5 Dai Z, Gao J, An G. 1996. Regulatory elements controlling developmental 6. and environmental regulation of a ribosomal protein L34 in tobacco. Plant Mol Biol In press.
- 7. Ausubel FM, Brent R, Kingson RE, Moore DD, Seidman JG, Smith JA, 10 Struhl K. 1992. Current protocols in molecular biology. John Wiley & Sons, New York.
- Marks JD, Hoogenboom HR, Griffiths AD, Winter G. 1992. Molecular 8. evolution of proteins on filamentous phage. Mimicking the strategy of the 15 immune system. J Biol Chem 267:16007-16010.
 - 9. An G. 1987. Binary Ti vectors for plant transformation and promoter analysis. Meth Enzy 153:292-305.
- Ishida Y, Saito H, Ohta S, Hiei Y, Komari T, Kumashiro T. 1996. High 20 efficiency transformation of maize (Zea mays L.) Mediated by Agrobacterium tumefaciens. Nature Biotechnol 14:745-750.
- Magnuson NS, Linzmaier PM, Gao J, Reeves R, An G, Lee JM. 1996. 25 Enhanced recovery of a secreted mammalian protein from suspension culture of genetically modified tobacco cells. Prot Expr Purif 7:220-228.
- Chrispeels MJ, Dickinson CD, Tague BW, Hunt DC, von Schaewen A. 1991. Defining the vacuolar targeting signal of phytohemagglutinin. In: 30 Plant Molecular Biology 2. Hermann RG, Larkins B, Eds. Plenum Press, New York. pp. 575-582.
- 13. Hellebust H, Uhlen M, Enfors SO. 1989. Effect of protein fusion on the stability of proteolytically sensitive sites in recombinant DNA proteins. J Biotechnol 12:275-284. 35
 - 14. Sijmons PC, Dekker BMM, Schrammeieijer B, Verwoerd TC, van den Elzen JM, Hoekema A. 1990. Production of correctly processed human serum albumin in transgenic plants. Bio/Technology 8:217-221.
 - 15. Verwoerd TC, van Paridon PA, van Ooyen AJJ, van Lent JWM, Hoekema A, Pen J. 1995. Stable accumulation of Aspergillus niger phytase in transgenic tobacco leaves. Plant Physiol 109:1199-1205.

- 16. Ellis JG, Llewellyn DJ, Walker JC, Dennis ES, Peacock WJ. 1987. The ocs element: a 16 base pair palindrome essential for activity of the octopine synthase enhancer. *EMBO J* 6:3203-3208.
- 5 17. Kay R, Chan A, Dayly M, McPherson J. 1987. Duplication of CaMV 35S promoter sequences creates a strong enhancer for plant genes. Science 236:1299-1302.
- Jobling SA, Gehrke L. 1987. Enhanced transcription of chimaeric
 messenger RNAs containing a plant viral untranslated leader sequence.
 Nature 325:622-625.
- 19. Cornelissen BJC, Hooft van Huysduynen RAM, Bol JF. 1986. A tobacco mosaic virus-induced tobacco protein is homologous to the sweet-tasting protein thaumatin. *Nature* 321:531-532.
 - 20. Hunt DC, Chrispeels MJ. 1991. The signal peptide of a vacuolar protein is necessary and sufficient for the efficient secretion of a cytosolic protein. *Plant Physiol* 96:18-25.
- Bell GI, Fong NM, Stempien MM, Wormsted MA, Caput D, Ku, L, Urdea MS, Rall LB, Sanchez-Pescador R. 1986. Human epidermal growth factor precursor: cDNA sequence, expression in vitro and gene. Nucleic Acids Res 14:8427-8446.
 - 22. Higo K, Saito Y, Higo H. 1993. Expression of a chemically synthesized gene for human epidermal growth factor under the control of cauliflower mosaic virus 35S promoter in transgenic tobacco. *Biosci Biotechnol Biochem* 57:1477-1481.

-16-

```
(1) GENERAL INFORMATION:
            APPLICANT: Brian S. Hooker, et al TITLE OF INVENTION: Method of Producing Human Growth
      Factors From Whole Plants or Plant Cell Cultures
      (iii) NUMBER OF SEQUENCES: 8
       (iv) CORRESPONDENCE ADDRESS:
                   (A) ADDRESSEE: Paul W. Zimmerman
                        STREET:
                   (B)
                                          P.O. Box 999
                   (C)
                        CITY:
                                      Richland
                   (D)
                        STATE:
                                          WA
                   (E)
                        COUNTRY:
                                          USA
                   (F)
                        ZIP:
                                          99352
             COMPUTER READABLE FORM:
       (v)
                   (A) MEDIUM TYPE: 3 1/2 Magnetic Disk
                   (B) COMPUTER: IBM compatible
                   (C) OPERATING SYSTEM: DOS
                   (D) SOFTWARE: WORD97
      (vi)
             CURRENT APPLICATION DATA:
                   (A) APPLICATION NUMBER: 08/747,246
                   (B) FILING DATE: 11-12-96
                   (C) CLASSIFICATION: unknown
      (vii) PRIOR APPLICATION DATA:
                   (A) APPLICATION NUMBER: N/A
                   (B) FILING DATE:
      (viii) ATTORNEY/AGENT INFORMATION:
                   (A) NAME: Paul W. Zimmerman (B) REGISTRATION NUMBER: 34,761
                   (C) REFERENCE/DOCKET NUMBER: E-1519
             TELECOMMUNICATION INFORMATION:
      (ix)
                   (A) TELEPHONE: 509-375-2981
(B) TELEFAX: 509-375-2592
                   (C) TELEX:
```

-17-

```
(2)
     INFORMATION FOR SEQ ID NO: 1:
      (i)
            SEQUENCE CHARACTERISTICS:
                 (A) LENGTH: 4481bp
                 (B) TYPE: Nucleic acid
                 (C) STRANDEDNESS: double strands
            (D) TOPOLOGY: unknown MOLECULE TYPE:
      (ii)
            (A) DESCRIPTION: Sense orientation of complementary DNA
                                 for pro-EGF
      (iii) HYPOTHETICAL:
      (iv) ANTI-SENSE: 5'-AGT GAC TCA GTC GAG ... TTC TCA CTC
                        GTC-3 end
            FRAGMENT TYPE:
      (v)
                              4.5kb Smal/Hindlll double strands DNA
                              fragment
      (vi)
            ORIGINAL SOURCE:
                 (A) ORGANISM: kidney
                 (B) STRAIN: human
                 (C) INDIVIDUAL ISOLATE: GI Belle
                 (D) DEVELOPMENTAL STAGE: adult
                 (E) HAPLOTYPE:
                 (F) TISSUE TYPE:
                 (G) CELL TYPE:
                 (H) CELL LINE:
                 (I) ORGANELLE:
      (vii) IMMEDIATE SOURCE:
                 (A) LIBRARY: fetal human liver library
                 (B) CLONE: lambda CH4A; lambda EMBL4; lambda GM1416
      (viii) POSITION IN GENOME:
                 (A) CHROMOSOME/SEGMENT:
                 (B) MAP POSITION:
                 (C) UNITS:
      (ix)
            FEATURE:
                 (A) NAME/KEY: human epithelial growth factor cDNA
                 (B) LOCATION:
                 (C) IDENTIFICATION METHOD: cross-hybridization
                                             with mouse cDNA
                 (D) OTHER INFORMATION:
            PUBLICATION INFORMATION:
      (x)
                 (A) AUTHORS:
                 (B) TITLE:
                 (C) JOURNAL:
                 (D) VOLUME:
                 (E) ISSUE:
                 (F)
                    PAGES:
                 (G) DATE:
                 (H) DOCUMENT NUMBER:
                 (I) FILING DATE:
                 (J) PUBLICATION DATE:
                    RELEVANT RESIDUES IN SEQ ID NO:
            FROM (position) TO SEQUENCE DESCRIPTION: SEQ ID NO: 1:
                                                   _ (position)
CCCGGGCCAT GCTCCAGCAA AATCAAGCTG TTTTCTTTTG AAAGTTCAAA CTCATCAAGA TT
ATG CTG CTC ACT CTT ATC ATT CTG TTG CCA GTA GTT TCA AAA TTT AGT TTT GTT 116
AGT CTC TCA GCA CCG CAG CAC TGG AGC TGT CCT GAA GGT ACT CTC GCA GGA AAT 170
GGG AAT TCT ACT TGT GTG GGT CCT GCA CCC TTC TTA ATT TTC TCC CAT GGA AAT
AGT ATC TTT AGG ATT GAC ACA GAA GGA ACC AAT TAT GAG CAA TTG GTG GTG GAT
GCT GGT GTC TCA GTG ATC ATG GAT TTT CAT TAT AAT GAG AAA AGA ATC TAT TGG 332
GTG GAT TTA GAA AGA CAA CTT TTG CAA AGA GTT TTT CTG AAT GGG TCA AGG CAA 386
GAG AGA GTA TGT AAT ATA GAG AAA AAT GTT TCT GGA ATG GCA ATA AAT TGG ATA 440
AAT GAA GAA GTT ATT TGG TCA AAT CAA CAG GAA GGA ATC ATT ACA GTA ACA GAT
                                                                          494
ATG AAA GGA AAT AAT TCC CAC ATT CTT TTA AGT GCT TTA AAA TAT CCT GCA AAT 548
GTA GCA GTT GAT CCA GTA GAA AGG TTT ATA TTT TGG TCT TCA GAG GTG GCT GGA 602
AGC CTT TAT AGA GCA GAT CTC GAT GGT GTG GGA GTG AAG GCT CTG TTG GAG ACA 656
```

```
TCA GAG AAA ATA ACA GCT GTG TCA TTG GAT GTG CTT GAT AAG CGG CTG TTT TGG 710
                AGA GAA
                        GGA AGC AAT
                                    TCT CTT ATT
                                                 TGC
            AAC
                                                     TCC
                                                         TGT
                                                              GAT
                                                                 TAT GAT 764
GGA GGT
        TCT
                        AGT
            GTC
                CAC
                    ATT
                            AAA
                                CAT CCA ACA
                                             CAG
                                                 CAT
                                                     AAT
                                                         TTG
                                                             TTT
                                                                  GCA
TCC CTT
        TTT GGT
                GAC
                    CGT
                        ATC
                            TTC
                                TAT TCA ACA TGG AAA ATG
                                                         AAG ACA ATT
                                                                      TGG 872
                                            AGA ATT
                                                     AAC CTC CAT TCA TCA 926
ATA GCC AAC AAA CAC ACT
                        GGA AAG GAC ATG GTT
                GGT GAA CTG AAA GTA GTG CAT
TTT GTA CCA CTT
                                             CCA CTT
                                                     GCA CAA CCC AAG GCA 980
        GAC
            ACT
                TGG
                    GAG
                        CCT
                            GAG CAG
                                    AAA CTT
   GAT
                                             TGC
                                                 AAA
                                                     TTG
                                                         AGG AAA
                                                                  GGA AAC
                                                                          1034
TGC AGC AGC ACT
                        GGG CAA GAC CTC CAG
                GTG
                    TGT
                                             TCA
                                                 CAC
                                                     TTG
                                                         TGC
                                                             ATG
                                                                  TGT
                                                                      GCA 1088
        TAC GCC
GAG GGA
                CTA
                    AGT
                        CGA
                            GAC CGG AAG TAC
                                             TGT
                                                 GAA GAT
                                                         GTT AAT
                                                                  GAA TGT 1142
GCT TTT
        TGG AAT
                CAT GGC
                        TGT ACT CTT GGG TGT
                                             AAA AAC ACC CCT GGA TCC
                                                                      TAT 1196
        ACG
            TGC
                CCT
                    GTA
                        GGA
                            TTT
                                GTT
                                    CTG CTT
                                             CCT
                                                 GAT
                                                     GGG
                                                         AAA CGA TGT CAT 1250
   CTT
CAA
        GTT
            TCC
                TGT
                    CCA CGC
                            AAT
                                GTG TCT GAA
                                             TGC
                                                 AGC
                                                     CAT
                                                         GAC
                                                             TGT
                                                                  GTT
                                                                      CTG
ACA TCA GAA GGT
                CCC
                    TTA
                        TGT
                            TTC TGT CCT GAA
                                             GGC
                                                 TCA GTG CTT
                                                             GAG AGA GAT 1358
GGG AAA
       ACA TGT
                AGC
                    GGT
                        TGT TCC TCA CCC GAT
                                            AAT
                                                 GGT GGA TGT AGC CAG CTC 1412
TGC GTT
        CCT CTT
                AGC
                    CCA
                        GTA
                            TCC
                                TGG GAA TGT
                                             GAT
                                                 TGC
                                                     TTT CCT GGG
                                                                 TAT GAC
CTA
   CAA
        CTG GAT
                GAA
                    AAA
                        AGC
                            TGT
                                GCA GCT
                                         TCA
                                             GGA
                                                 CCA
                                                     CAA CCA
                                                             TTT
                                                                  TTG
                                                                      CTG
TTT GCC
       AAT TCT
                CAA GAT
                        ATT
                            CGA CAC ATG CAT
                                             TTT
                                                 GAT GGA ACA GAC TAT
                                                                      GGA 1574
ACT CTG
        CTC AGC
                CAG
                    CAG ATG GGA ATG GTT TAT GCC
                                                 CTA GAT CAT GAC CCT GTG 1628
GAA AAT
        AAG ATA
                TAC
                    TTT
                        GCC CAT ACA GCC CTG
                                            AAG
                                                 TGG ATA GAG AGA GCT AAT 1682
                    CGA GAA AGG CTT ATT GAG
ATG GAT
        GGT
            TCC
                CAG
                                             GAA
                                                 GGA GTA GAT
                                                             GTG CCA GAA
GGT CTT
        GCT GTG
                GAC
                    TGG ATT
                            GGC CGT AGA TTC
                                             TAT
                                                 TGG ACA GAC AGA GGG AAA 1790
                    AGT GAT TTA AAT GGG AAA
                                                 TCC AAA ATA ATC ACT AAG 1844
TCT CTG ATT GGA
                AGG
                                             CGT
GAG
   AAC
        ATC
            TCT
                CAA
                    CCA
                        CGA GGA ATT
                                    GCT GTT
                                             CAT
                                                 CCA ATG
                                                         GCC AAG AGA TTA 1898
TTC
    TGG
       ACT
            GAT
                ACA
                    GGG ATT AAT
                                CCA CGA ATT
                                             GAA
                                                 AGT
                                                     TCT
                                                         TCC
                                                             CTC
                                                                 CAA GGC
CTT GGC CGT
           CTG
                GTT ATA GCC AGC TCT GAT CTA ATC
                                                 TGG CCC AGT
                                                             GGA ATA ACG 2006
ATT GAC TTC TTA
                ACT GAC
                        AAG TTG TAC TGG TGC GAT
                                                 GCC AAG CAG TCT GTG ATT 2060
                                                         CAG AAT GAT GTA 2114
GAA ATG
        GCC AAT
                CTG GAT GGT TCA AAA CGC CGA AGA CTT ACC
    CAC
        CCA
            TTT
                GCT
                    GTA GCA GTG TTT
                                    GAG GAT
                                             TAT
                                                 GTG
                                                     TGG
                                                         TTC
                                                             TCA GAT
                                                                      TGG
GCT ATG CCA TCA
                GTA ATA AGA GTA AAC AAG AGG ACT
                                                 GGC AAA GAT AGA GTA CGT 2222
CTC CAA GGC AGC
                ATG CTG AAG CCC TCA TCA CTG
                                            GTT
                                                 GTG GTT CAT CCA TTG GCA 2276
                        TGC TTA TAT CAA AAC GGA
AAA CCA GGA GCA
                GAT
                    CCC
                                                 GGC
                                                     TGT
                                                         GAA CAT ATT TGC 2330
                            TGG
                                                 GAA GGT
                                                         TTT
AAA AAG
        AGG
            CTT
                GGA
                    ACT
                        GCT
                                TGT
                                    TCG
                                        TGT
                                             CGT
                                                             ATG AAA GCC 2384
                                                                      GGT 2438
TCA GAT
        GGG AAA
                ACG
                    TGT
                        CTG GCT
                                CTG GAT
                                         GGT
                                             CAT
                                                 CAG CTG TTG GCA GGT
GAA GTT GAT CTA AAG AAC
                        CAA GTA ACA CCA TTG GAC ATC
                                                     TTG TCC AAG ACT AGA 2492
GTG TCA GAA GAT AAC ATT ACA GAA TCT CAA CAC
                                            ATG CTA GTG GCT GAA ATC ATG 2546
GTG
    TCA GAT CAA GAT
                    GAC
                        TGT GCT CCT GTG GGA
                                             TGC AGC ATG
                                                         TAT GCT CGG TGT 2600
                                        TGT
TTA
    TCA GAG
            GGA
                GAG
                    GAT
                        GCC ACA TGT CAG
                                             TTG
                                                 AAA GGA
                                                         TTT GCT
                                                                  GGG GAT
GGA AAA CTA
            TGT
                TCT
                    GAT ATA GAT GAA TGT GAG ATG GGT GTC CCA GTG TGC CCC 2708
CCT GCC
        TCC TCC
                AAG
                    TGC
                        ATC AAC ACC GAA GGT
                                             GGT TAT GTC
                                                         TGC CGG TGC TCA 2762
                        GGG ATT CAC
                                    TGT CTT
                                                             TGC CAA CTG 2816
GAA GGC
        TAC CAA
                GGA GAT
                                             GAT ATT
                                                     GAT GAG
GGG
    GTG
        CAC
            AGC
                TGT
                    GGA
                        GAG AAT
                                 GCC
                                    AGC
                                        TGC
                                             ACA AAT
                                                     ACA
                                                         GAG
                                                             GGA
                                                                  GGC
                                                                      TAT
                    GGA CGC CTG
                                TCT GAA CCA GGA AAT AGT GAC
                                                                      TGT 2924
ACC
    TGC ATG TGT
                GCT
                                                             TCT
                                                                 GAA
CCC CTG TCC CAC
                GAT
                    GGG
                        TAC TGC
                                CTC CAT GAT
                                             GGT GTG TGC ATG
                                                             TAT ATT GAA 2978
   TTG GAC AAG TAT
GCA
                    GCA TGC AAC
                                TGT GTT GTT GGC
                                                 TAC ATC GGG GAG CGA TGT 3032
                                                                  TCT
                    AAG
                        TGG
                            TGG
                                 GAA CTG CGC
                                             CTG
                                                 ATT
                                                     TGC
                                                         CCT
                                                             GAC
    TAC
        CGA
            GAC
                CTG
                                                                      ACT
        CCT
                    AGG
                        GAA GAT
                                GAC CAC CAC
                                             TAT
                                                 TCC
                                                     GTA AGA CAC
                                                                  GCT
                                                                          3140
CCA CCC
            CAC
                CTC
                                                                      GGC
CAC GGG CAG CAG
                CAG
                    AAG
                        GTC ATC GTG GTG GCT
                                             GTC TGC
                                                     GTG GTG
                                                             GTG CTT
                                                                      GTC 3194
ATG CTG CTC CTC
                    AGC
                        CTG TGG GGG GCC CAC
                                             TAC
                                                 TAC AGG ACT
                                                             CAG AAG CTG 3248
                CTG
CTA
    TCG AAA AAC
                    AAG
                        AAT
                            CCT
                                 TAT
                                    GAG
                                         GAG
                                             TCG
                                                 AGC
                                                     AGA
                                                         GAT
                                                             GTG
                CCA
                                                                  AGG
                                                                      AGT
                                 GGG ATG TCC
                                                                  TGG TTT 3356
CGC AGG CCT GCT
                GAC ACT
                        GAG GAT
                                             TCT
                                                 TGC CCT CAA CCT
GTG GTT ATA AAA
                GAA CAC CAA GAC CTC AAG AAT GGG GGT CAA CCA GTG GCT GGT 3410
GAG GAT GGC CAG GCA GCA GAT
                            GGG TCA ATG CAA CCA ACT
                                                     TCA TGG AGG
                                                                 CAG GAG
                                                                          3464
                        GGC
                            ACA GAG CAA GGC
                                             TGC
                                                 TGG
                                                     ATT
                                                         CCA
                                                             GTA
        TTA
            TGT
                GGA ATG
                                                                  TCC
                                                                      AGT
GAT AAG GGC TCC
                TGT CCC
                        CAG GTA ATG GAG CGA AGC
                                                 TTT CAT
                                                         ATG CCC
                                                                  TCC
                                                                      TAT
                                                                          3572
GGG ACA CAG ACC CTT GAA GGG GGT GTC GAG AAG CCC CAT TCT CTC CTA TCA GCT 3626
AAC CCA TTA TGG CAA CAA AGG GCC CTG GAC CCA CCA CAC CAA ATG GAG CTG ACT 3680
                                                                          3746
AAACTGGAAT TAAAAGGAAA GTCAAGAAGA ATGAACTATG TCGATGCACA GTATCTTTTC
TTTCAAAAGT AGAGCAAAAC TATAGGTTTT GGTTCCACAA TCTCTACGAC TAATCACCTA
                                                                          3806
CTCAATGCCT GGAGACAGAT ACGTAGTTGT GCTTTTGTTT GCTCTTTTAA GCAGTCTCAC
                                                                          3866
TGCAGTCTTA TTTCCAAGTA AGAGTACTGG GAGAATCACT AGGTAACTTA TTAGAAACCC
AAATTGGGAC AACAGTGCTT TGTAAATTGT GTTGTCTTCA GCAGTCAATA CAAATAGATT
                                                                          3986
TTTGTTTTTG TTGTTCCTGC AGCCCCAGAA GAAATTAGGG GTTAAAGCAG ACAGTCACAC
TGGTTTGGTC AGTTACAAAG TAATTTCTTT GATCTGGACA GAACATTTAT ATCAGTTTCA
                                                                          4106
TGAAATGATT GGAATATTAC AATACCGTTA AGATACAGTG TAGGCATTTA ACTCCTCATT
GGCGTGGTCC ATGCTGATGA TTTTGCCAAA ATGAGTTGTG ATGAATCAAT GAAAAATGTA
                                                                          4226
```

-19-

ATTTAGAAAC TGATTTCTTC AGAATTAGA ACATTTTATT TTTAAAATAT TACACAGGA TTTTCCCCTA CAGAATTTTC CCTCTTGGT TTACAAGATT GTAAGTAAAT TGCCTGATT CTAATTATGA ATTC	GCCTTCGGAG GTGATTGCACA	TTTCTTAGTC GAATTTGTAT	ATTACTGTCC GTATTTTCAG	4286 4346 4406 4466 4480
---	---------------------------	--------------------------	--------------------------	--------------------------------------

```
INFORMATION FOR SEQ ID NO: 2:
             SEQUENCE CHARACTERISTICS:
                   (A) LENGTH: 783bp
                   (B) TYPE: Nucleic acid
                   (C) STRANDEDNESS: double strands
                   (D) TOPOLOGY: unknown
             MOLECULE TYPE:
       (ii)
                   (A) DESCRIPTION: sense orientation of five
                                     copies of mature EGF concatemers
       (iii) HYPOTHETICAL:
       (iv) ANTI-SENSE: 5'-CGC GTC AAG GGT ... TCT CAG TGA TAA-3
                                end
             FRAGMENT TYPE: 4.5kb Smal/Hindlll double strands DNA
       (v)
                                fragment
             ORIGINAL SOURCE:
                   (A) ORGANISM: kidney
                   (B) STRAIN: human
                   (C) INDIVIDUAL ISOLATE: Z.Dai, et al.
                   (D) DEVELOPMENTAL STAGE: adult
                  (E) HAPLOTYPE:
                   (F) TISSUE TYPE:
                  (G) CELL TYPE:
                  (H) CELL LINE:
                  (I) ORGANELLE:
       (vii) IMMEDIATE SOURCE:
                  (A) LIBRARY: fetal human liver library
                  (B) CLONE: lambda CH4A; lambda EMBL4; lambda GM1416
       (viii) POSITION IN GENOME:
                  (A) CHROMOSOME/SEGMENT:
                  (B) MAP POSITION:
                  (C) UNITS:
             FEATURE:
       (ix)
                  (A) NAME/KEY: Concatemer of mature EGF fragment
                                 without linker
                  (B) LOCATION:
                  (C) IDENTIFICATION METHOD: PCR cloning
                  (D) OTHER INFORMATION:
       (x)
             PUBLICATION INFORMATION:
                  (A) AUTHORS:
                  (B) TITLE:
                  (C) JOURNAL:
                  (D) VOLUME:
                  (E) ISSUE:
                  (F) PAGES:
                  (G) DATE:
                  (H) DOCUMENT NUMBER:
                  (I) FILING DATE:
                  (J) PUBLICATION DATE:
                  (K) RELEVANT RESIDUES IN SEQ ID NO:
            FROM (position) TO (position) SEQUENCE DESCRIPTION: SEQ ID NO: \overline{2}:
      (xi)
AAT AGT GAC TCT GAA TGT CCC CTG TCC CAC GAT GGG TAC TGC CTC CAT GAT GGT 54
GTG TGC ATG TAT ATT GAA GCA TTG GAC AAG TAT GCA TGC AAC TGT GTT GGC 108
TAC ATC GGG GAG CGA TGT CAG TAC CGA GAC CTG AAG TGG TGG GAA CTG CGC AAT 162 AGT GAC TCT GAA TGT CCC CTG TCC CAC GAT GGG TAC TGC CTC CAT GAT GGT GTG 216
TGC ATG TAT ATT GAA GCA TTG GAC AAG TAT GCA TGC AAC TGT GTT GGC
                                                                         TAC
ATC GGG GAG CGA TGT CAG TAC CGA GAC CTG AAG TGG TGG GAA CTG CGC AAT AGT
    TCT GAA TGT CCC CTG TCC CAC GAT GGG TAC TGC CTC CAT GAT GGT GTG TGC 378
ATG TAT ATT GAA GCA TTG GAC AAG TAT GCA TGC AAC TGT GTT GGC TAC ATC 432
```

SUBSTITUTE SHEET (RULE 26)

GGG GAG CGA TGT CAG TAC CGA GAC CTG AAG TGG TGG GAA CTG CGC AAT AGT GAC 486

WO 98/21348 PCT/US97/20603

-21-

TCT	GAA	TGT	CCC	CTG	TCC	CAC	GAT	GGG	TAC	TGC	CTC	CAT	GAT	GGT	GTG	TGC	ATG	540
TAT	ATT	GAA	GCA	TTG	GAC	AAG	TAT	GCA	TGC	AAC	TGT	GTT	GTT	GGC	TAC	ATC	GGG	594
GAG	CGA	TGT	CAG	TAC	CGA	GAC	CTG	AAG	TGG	TGG	GAA	CTG	CGC	AAT	AGT	GAC	TCT	648
GAA	TGT	CCC	CTG	TCC	CAC	GAT	GGG	TAC	TGC	CTC	CAT	GAT	GGT	GTG	TGC	ATG	TAT	702
ATT	GAA	GCA	TTG	GAC	AAG	TAT	GCA	TGC	AAC	TGT	GTT	GTT	GGC	TAC	ATC	GGG	GAG	756
CGA	TGT	CAG	TAC	CGA	GAC	CTG	AAG	TGG	TGG	GAA	CTG	CGC					•	795

```
INFORMATION FOR SEQ ID NO: 3:
(4)
               SEQUENCE CHARACTERISTICS:
                      (A) LENGTH: 891bp
                      (B) TYPE: Nucleic acid
                      (C) STRANDEDNESS: double strands
               (D) TOPOLOGY: unknown MOLECULE TYPE:
        (ii)
                      (A) DESCRIPTION: sense orientation concatemer of
                                            mature EGF fragments with linkers
        (iii) HYPOTHETICAL:
        (iv) ANTI-SENSE: 5'-GTC CAG AGC ... CAG TGA TAA-3 end
(v) FRAGMENT TYPE: 5-copies of 159bp concatemer mature EGF
                                  linked with linkers
               ORIGINAL SOURCE:
        (vi)
                      (A) ORGANISM: kidney
                      (B) STRAIN: human
                      (C) INDIVIDUAL ISOLATE: Z. Dai, et al
                      (D) DEVELOPMENTAL STAGE: adult
                      (E) HAPLOTYPE:
                      (F) TISSUE TYPE:
                      (G) CELL TYPE:
                      (H) CELL LINE:
                      (I) ORGANELLE:
        (vii) IMMEDIATE SOURCE:
                      (A) LIBRARY:
(B) CLONE:
        (viii) POSITION IN GENOME:
                      (A) CHROMOSOME/SEGMENT:
                      (B) MAP POSITION:
                      (C) UNITS:
               FEATURE:
        (ix)
                      (A) NAME/KEY: concatemer of mature EGF linked with linkers
                      (B) LOCATION:
                      (C) IDENTIFICATION METHOD: PCR cloning
                      (D) OTHER INFORMATION: Cleavage sites at 142-165, 307-331,
465-489, 631-655.
               PUBLICATION INFORMATION:
        (x)
                      (A) AUTHORS:
                      (B) TITLE:
                      (C) JOURNAL:
                      (D) VOLUME:
(E) ISSUE:
(F) PAGES:
                      (G) DATE:
                      (H) DOCUMENT NUMBER:
                      (I) FILING DATE:(J) PUBLICATION DATE:
                      (K) RELEVANT RESIDUES IN SEQ ID NO:
               FROM (position) TO (position) SEQUENCE DESCRIPTION: SEQ ID NO: 3:
        (xi)
AAT AGT GAC TCT GAA TGT CCC CTG TCC CAC GAT GGG TAC TGC CTC CAT GAT GGT 54 GTG TGC ATG TAT ATT GAA GCA TTG GAC AAG TAT GCA TGC AAC TGT GTT GTT GGC 10 TAC ATC GGG GAG CGA TGT CAG TAC CGA GAC CTG AAG TGG TGG GAA CTG CGC GGC 16 GGA AGA GTT AAC TGC ATG CAG AAT AGT GAC TCT GAA TGT CCC CTG TCC CAC GAT 21
                                                                                            108
                                                                                            162
                    TGC ATG CAG AAT AGT CAT GAT GGT GTG TGC
                                                                                            216
               CTC
                    CAT
                         GAT GGT
                                              ATG TAT ATT GAA GCA
GGG TAC
          TGC
                                                                       TTG GAC
                                                                                 AAG
                                                                                       TAT
GCA TGC
               TGT
                         GTT GGC TAC ATC GGG GAG CGA TGT CAG TAC
          AAC
                    GTT
                                                                            CGA
                                                                                 GAC
                                                                                       CTG
                         CGC GGC GGA AGA CAC GAT GGG TAC
                                              GTT AAC
TGC CTC
AAG TGG TGG
                    CTG
                                                        TGC ATG CAG AAT
                                                                            AGT
                                                                                 GAC
               GAA
                                                                                       TCT
                                                                                            378
                                                             GAT
GTT
GAA TGT
                                              TGC
          CCC
               CTG
                    TCC
                                                        CAT
                                                                  GGT
                                                                       GTG
                                                                            TGC
                                                                                 ATG
                                                                                       TAT
ATT GAA
          GCA
               TTG
                    GAC
                         AAG TAT
                                   GCA TGC
                                              AAC TGT
                                                        GTT
                                                                  GGC
                                                                       TAC
                                                                            ATC
                                                                                 GGG
                                                                                       GAG
                                                                                            486
                    CGA GAC CTG
AGT GAC TCT
CGA TGT
          CAG
                              CTG AAG TGG
                                              TGG GAA
                                                        CTG CGC
                                                                  GGC
                                                                       GGA AGA
               TAC
                                                                                 GTT
                                                                                       AAC
                                                                                            540
TGC ATG
          CAG AAT
                    AGT
                                   GAA TGT
                                              CCC CTG
                                                        TCC CAC
                                                                  GAT
                                                                       GGG
                                                                            TAC
                                                                                 TGC
                                                                                       CTC
CAT GAT
                         ATG TAT ATT
                                        GAA GCA TTG
                                                        GAC AAG
                                                                  TAT
                                                                       GCA
                                                                                            648
                                                                            TGC
                                                                                 AAC
          GGT
                                                                                       TGT
               GTG
                    TGC
GTT GTT
                                              CAG TAC
                                                                  CTG
                                                                       AAG
                                                                                            702
                                   CGA
                                        TGT
                                                             GAC
                                                                            TGG
          GGC
               TAC
                    ATC
                         GGG
                              GAG
                                                        CGA
                                                                                 TGG
                                                                                       GAA
                         GTT AAC TGC ATG
TGC CTC CAT GAT
                                   TGC ATG CAG AAT AGT GAC
                                                                  TCT GAA
CTG CGC
          GGC
               GGA AGA
                                                                            TGT CCC
                                                                                       CTG
                                                                                            756
                                              GGT GTG TGC ATG TAT ATT GAA
GGC TAC ATC GGG GAG CGA TGT
TCC
     CAC
          GAT
               GGG
                    TAC
                                                                                  GCA
                                                                                       TTG
                                                                                            810
          TAT GCA
                    TGC
                         AAC TGT GTT GTT
GAC AAG
```

CGA GAC CTG AAG TGG TGG GAA CTG CGC

-23-

```
INFORMATION FOR SEQ ID NO: 4:
(5)
       (i.)
            SEQUENCE CHARACTERISTICS:
                    (A) LENGTH: 330bp
(B) TYPE: nucleic acid
                    (C) STRANDEDNESS: double strands
                    (D) TOPOLOGY: unknown
       (ii) MOLECULE TYPE:
                    (A) DESCRIPTION: upstream enhancer (from -343 to
                                         -90 bp) of 35S promoter
       (iii) HYPOTHETICAL:
       (iv) ANTI-SENSE:
             FRAGMENT TYPE: 253bp upstream of 35S promoter
                               enhancer element
       (vi) ORIGINAL SOURCE:
                    (A) ORGANISM: cauliflower mosaic virus (CaMV)
                    (B) STRAIN: Cabb B-D
                    (C) INDIVIDUAL ISOLATE: Z.Dai, et al
                    (D) DEVELOPMENTAL STAGE:
                    (E) HAPLOTYPE:
                    (F) TISSUE TYPE:
                    (G) CELL TYPE:
                    (H) CELL LINE:
                    (I) ORGANELLE:
       (vii) IMMEDIATE SOURCE:
                    (A) LIBRARY:
                    (B) CLONE:
       (viii) POSITION IN GENOME:
                    (A) CHROMOSOME/SEGMENT:
                    (B) MAP POSITION:
(C) UNITS:
       (ix) FEATURE:
                    (A) NAME/KEY: 35S promoter B-domain enhancer
                    (B) LOCATION:
(C) IDENTIFICATION METHOD: standard cloning
                    (D) OTHER INFORMATION: B-domain of 35S promoter from EcoR V
site to Hind II site (upstream enhancer region from -343 to -90 bp)
             PUBLICATION INFORMATION:
                    (A) AUTHORS:
                    (B) TITLE:
                    (C) JOURNAL:
                    (D) VOLUME:
(E) ISSUE:
                    (F) PAGES:
                    (G) DATE:
                    (H) DOCUMENT NUMBER:
                    (I) FILING DATE:
                    (J) PUBLICATION DATE:
                    (K) RELEVANT RESIDUES IN SEQ ID NO:
       FROM (position) TO (position) (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 4:
GTCAACATGG TGGAGCACGA CACACTTGTC TACTCCAAAA ATATCAAAGA TACAGTCTCA 60
GAAGACCAAA GGGCAATTGA GACTTTTCAA CAAAGGGTAA TATCCGGAAA CCTCCTCGGA 120
TTCCATTGCC CAGCTATCTG TCACTTTATT GTGAAGATAG TGGAAAAGGA AGGTGGCTCC 180 TACAAATGCC ATCATTGCGA TAAAGGAAAG GCCATCGTTG AAGATGCCTC TGCCGACAGT 240
GGTCCCAAAG ATGGACCCCC ACCCACGAGG AGCATCGTGG AAAAAGAAGA CGTTCCAACC 300
```

ACGTCTTCAA AGCAAGTGGA TTGATGTGAT

```
INFORMATION FOR SEO ID NO: 5:
              SEQUENCE CHARACTERISTICS:
       (i)
                    (A) LENGTH: 1441bp
                    (B) TYPE: Nucleic acid
                    (C) STRANDEDNESS: double strands
                    (D) TOPOLOGY: unknown
              MOLECULE TYPE:
       (ii)
                    (A) DESCRIPTION: 5'-untranscription region of
                                        chl a/b binding protein
       (iii) HYPOTHETICAL:
       (iv) ANTI-SENSE:
              FRAGMENT TYPE: 11kb EcoR 1 fragment
       (v)
       (vi)
              ORIGINAL SOURCE:
                    (A) ORGANISM: whole plants
                    (B) STRAIN: Arabidopsis
                    (C) INDIVIDUAL ISOLATE: Ha et al
                    (D) DEVELOPMENTAL STAGE: 30 day old seedlings
                    (E) HAPLOTYPE:
                    (F) TISSUE TYPE:
                    (G) CELL TYPE:
                    (H) CELL LINE:
                    (I) ORGANELLE:
       (vii) IMMEDIATE SOURCE:
                    (A) LIBRARY: genomic DNA library
                    (B) CLONE: lambda bAT1005
       (viii) POSITION IN GENOME:
                    (A) CHROMOSOME/SEGMENT:
                    (B) MAP POSITION:
                    (C) UNITS:
              FEATURE:
       (ix)
                    (A) NAME/KEY: arabidopsis cabl gene promoter
                    (B) LOCATION:
                    (C) IDENTIFICATION METHOD: cross-hybridization
                    (D) OTHER INFORMATION:
              PUBLICATION INFORMATION:
       (x)
                    (A) AUTHORS:
                    (B) TITLE:
                    (C) JOURNAL:
                    (D) VOLUME:
(E) ISSUE:
                    (F) PAGES:
                    (G) DATE:
                    (H) DOCUMENT NUMBER:
                    (I) FILING DATE:
                    (J) PUBLICATION DATE:
                    (K) RELEVANT RESIDUES IN SEQ ID NO:
       FROM (position) TO (position) (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 5:
GAATTCATCA ACAAATTACT CCTCAATCAC ACTCCTATAG AAAACGGTTT AAGCTATCAT 60
TACATGTCTA GTTGGTTTTA CTCAGCCCTA GAAGTGTTGT TTATTGCATC ACTTTCCACG 120
AAGCACAATT TTTCTTTTTT ACAATCACTA GACCTCACAG GCTCACACAT ATGCTTTAGA 180
GCACATTCTA AACTTTGAAC TATAAAAGCT GTTAACACTA ATACACTATG CGTTCTTTTT TGCTCCAAAC ACTTTTGATC CATTATTAGG AGACACTCCA CTTAGAAAGA TTTTCTAATC
                                                                              240
                                                                              300
CTTTGGTCAA CTAGGAAGTT CAAGGTTTTT CTAAACAGAA ATTCATTTCA CAAGTAATTT 360
AATTTATAAG GAAATGAATA GAGAAATCAA ATCATTGAAG AACTACAAAA TATAGATTCA 420
AGGTCAGGTC TAAGAAAATA TTCCTGAAGC TCAAAAAAGA GTTTTCCTCT CACATTATAG 480
AATTGGCCTT TACTTCAACA TTTTCCCACC TATTCCACAT TTGGTCAGAA CATTTTTAAT 540
TACTTGTGGA TCAATTTCCG GTTGAAATGG GTTTGGTGAA TATCCGGTTC AGTTATATGG 600
TGGCCGTTGG AATTGGCTTA TTAGTTGTGG CCGTTGTTGA AGCCGTTGGT ATTGGTAAGG 660
GAGAAGCAGA CTTGTGGCTA TGAGTCTATG ACCATGACTC GTGATTATGG AGCTGTCTTA 720 TGACCCTGAC CATCACCTTG ATCTGGTGGA TTCCAATGTT TTCTTCTTCT TCTAATAAAA 780
TATTATGGTC AATACAGGTG CTAATTAAGA TGGTAATAAT TTCTTATGTT TCTGTGGTAA 840
AGTTTGATTC AATTCCGTAG TTTTAGATAA TCTTATTTCC ATACATAAAT TTTATAGTTT 900
TATCTACTTT GTTCTTATGT TTTATCTCTA GCCAAGAGTT ATTATTATTA TCAGAAGAAG 960
AAAAAAAAA GAAGCATATA TACAAAAGGT TTAATAAAAT GTATTATACA AGGCAATTAT 1020
```

WO 98/21348 PCT/US97/20603

-25-

CCAAATTTTT	TTTGTTTTGG	TTTACATTGA	TGCTCTCAGG	ATTTCATAAG	GATAGAGAGA	1080
TCTATTCGTA	TACGTGTCAC	GTCATGAGTG	GGTGTTTCGC	CAATCCATGA	AACGCACCTA	1140
GATATCTAAA	ACACATATCA	ATTGCGAATC	TGCGAAGTGC	GAGCCATTAA	CCACGTAAGC	1200
AAACAAACAA	TCTAAACCCC	AAAAAAAATC	TATGACTAGC	CAATAGCAAC	CTCAGAGATT	1260
GATATTTCAA	GATAAGACAG	TATTTAGATT	TCTGTATTAT	ATATAGCGAA	AATCGCATCA	1320
ATACCAAACC	ACCCATTTCT	TGGCTTACAA	CAACAAATCT	TAAACGTTTT	ACTTTGTGCT	1380
GCACTACTCA	ACCTTAATGG	CCGCCTCAAC	AATGGCTCTC	TCCTCCCCTG	CCTTCGCCGG	1440
T						1441

```
INFORMATION FOR SEQ ID NO: 6:
               SEQUENCE CHARACTERISTICS:
        (i)
                       (A) LENGTH: 832bp
                       (B) TYPE: Nucleic acid
                       (C) STRANDEDNESS: double strands
                       (D) TOPOLOGY: unknown
              MOLECULE TYPE:
                       (A) DESCRIPTION: CaMV 35S 5'-untranscription
                                             upstream
        (iii) HYPOTHETICAL:
        (iv) ANTI-SENSE:
        (v)
                FRAGMENT TYPE: Alu 1 (from 7143bp)-EcoR1(to 7517bp)
        (vi)
               ORIGINAL SOURCE:
                       (A) ORGANISM: cauliflower mosaic virus
                       (B) STRAIN: cM4-184
                       (C) INDIVIDUAL ISOLATE: RJ Shepherd
                       (D) DEVELOPMENTAL STAGE:
                       (E) HAPLOTYPE:
                       (F) TISSUE TYPE:
                       (G) CELL TYPE:
                       (H) CELL LINE:
                       (I) ORGANELLE:
        (vii) IMMEDIATE SOURCE:
                       (A) LIBRARY: genomic library of CM4-184
                       (B) CLONE: pOS-1
        (viii) POSITION IN GENOME:
                       (A) CHROMOSOME/SEGMENT:
                       (B) MAP POSITION:
                       (C) UNITS:
               FEATURE:
        (ix)
                       (A) NAME/KEY: CaMV 35S promoter
                       (B) LOCATION:
                       (C) IDENTIFICATION METHOD: cross-hybridization
                       (D) OTHER INFORMATION:
               PUBLICATION INFORMATION:
        (x)
                       (A) AUTHORS:
                       (B) TITLE:
                       (C) JOURNAL:
                       (D) VOLUME:
                       (E) ISSUE:
                      (F) PAGES: (G) DATE:
                       (H) DOCUMENT NUMBER:
                       (I) FILING DATE:
                       (J) PUBLICATION DATE:
                       (K) RELEVANT RESIDUES IN SEQ ID NO:
        FROM (position) TO ___ (position) (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 6:
CCCACA GATGGTTAGA GAGGCTTACG CAGCAGGTCT CATCAAGACG ATCTACCCGA 56
GCAATAATCT CCAGGAAATC AAATACCTTC CCAAGAAGGT TAAAGATGCA GTCAAAAGAT 116
TCAGGACTAA CTGCATCAAG AACACAGAGA AAGATATATT TCTCAAGATC AGAAGTACTA 176
TTCCAGTATG GACGATTCAA GGCTTGCTTC ACAAACCAAG GCAAGTAATA GAGATTGGAG 236
TCTCTAAAAA GGTAGTTCCC ACTGAATCAA AGGCCATGGA GTCAAAGATT CAAATAGAGG 296
ACCTAACAGA ACTCGCCGTA AAGACTGGCG AACAGTTCAT ACAGAGTCTC TTACGACTCA 356
ATGACAAGAA GAAAATCTTC GTCAACATGG TGGAGCACGA CACACTTGTC TACTCCAAAA 416
ATATCAAAGA TACAGTCTCA GAAGACCAAA GGGCAATTGA GACTTTTCAA CAAAGGGTAA 476
TATCCGGAAA CCTCCTCGGA TTCCATTGCC CAGCTATCTG TCACTTTATT GTGAAGATAG 536
TGGAAAAGGA AGGTGGCTCC TACAAATGCC ATCATTGCGA TAAAGGAAAG GCCATCGTTG 596
AAGATGCCTC TGCCGACAGT GGTCCCAAAG ATGGACCCCC ACCCACGAGG AGCATCGTGG 656
AAAAAGAAGA CGTTCCAACC ACGTCTTCAA AGCAAGTGGA TTGATGTGAT ATCTCCACTG 716
ACGTAAGGGA TGACGCACAA TCCCACTATC CTTCGCAAGA CCCTTCCTCT ATATAAGGAA 776
```

GTTCATTTCA TTTGGAGAGA ACACGGGGGA CTCTAGAGGA TCCCCGGGTG GTCAGT

```
(8)
      INFORMATION FOR SEQ ID NO: 7:
                SEQUENCE CHARACTERISTICS:
        (i)
                      (A) LENGTH: 473bp
                      (B) TYPE: Nucleic acid
(C) STRANDEDNESS: double strands
                      (D) TOPOLOGY: unknown
        (ii)
               MOLECULE TYPE:
                      (A) DESCRIPTION: 5'-untranscription upstream of
                                             ribosomal protein L34
        (iii) HYPOTHETICAL:
        (iv) ANTI-SENSE:
        (v)
                FRAGMENT TYPE: 1500bp BamH-Hind 111
        (vi)
               ORIGINAL SOURCE:
                      (A) ORGANISM: tobacco NT1 cells
                      (B) STRAIN:
                      (C) INDIVIDUAL ISOLATE: Z.Dai, et al
                      (D) DEVELOPMENTAL STAGE: 3 days old (E) HAPLOTYPE:
                      (F) TISSUE TYPE:
                      (G) CELL TYPE:
                      (H) CELL LINE: NT1
                      (I) ORGANELLE:
        (vii) IMMEDIATE SOURCE:
                      (A) LIBRARY: genomic library(B) CLONE: TSC 40
        (viii) POSITION IN GENOME:
                      (A) CHROMOSOME/SEGMENT:
                      (B) MAP POSITION:
                      (C) UNITS:
        (ix)
               FEATURE:
                      (A) NAME/KEY: RPL-34 promoter
                      (B) LOCATION:
                      (C) IDENTIFICATION METHOD: plaque hybridization (D) OTHER INFORMATION:
                PUBLICATION INFORMATION:
        (x)
                      (A) AUTHORS:
                      (B) TITLE:
                      (C) JOURNAL: (D) VOLUME:
                      (E) ISSUE:
                      (F) PAGES: (G) DATE:
                      (H) DOCUMENT NUMBER:
                      (I) FILING DATE:
                      (J) PUBLICATION DATE:
                      (K) RELEVANT RESIDUES IN SEQ ID NO:
        FROM (position) TO (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 7:
                                                            ____ (position)
AGATCTCT CTTTGTATTC TTATTGATGT ACTGGTTTGA AGATGAATAA AATCTTTCAT 58
TCCACCAAAA AAAGAATGAA AATAAAATTT TAATATACAT GTTGATATAG ACAAAGAAGA 118
AAAAAAAAGT TGTGATTACA TTTATTGACT ATTTGATGCC AATATCTATA ACTAGAGCTA 178
TTTTCTATCA ATTATATGGG TATGTTGTTA TACCATGCCA AAACCTCAAT TCATAATGTG 238
CTTGTTTAAA CCCAGTTTAA TGGGCTAACA TGTTGATGGG CTTATAGGCC CGTCTGATTT 298
CCTTGCCAGA CACTAGTAAG TAAATGATTC TATCATCCAA TATCAACCGT GGGATCTAGG 358
GCTTGTCCCA CTTATATACA CTACATATAT TTAACTTTCC TTTAGCCCTT CTGCTTCAGC 418
```

CCCCAAAACA AAGAAGAAG CTACAGAGAG AATAGCAGCG CCGCCGTGAA AAATG

-28-

```
INFORMATION FOR SEQ ID NO: 8:
(9)
               SEQUENCE CHARACTERISTICS:
                      (A) LENGTH: 1162bp
                      (B) TYPE: Nucleic acid
                      (C) STRANDEDNESS: double strands
                      (D) TOPOLOGY: unknown
        (ii)
               MOLECULE TYPE:
               (A) DESCRIPTION: 5'-untranscription region of 35S gene
                                     from CaMV with 2 copies of B domains
        (iii) HYPOTHETICAL:
        (iv) ANTI-SENSE:
        (v)
               FRAGMENT TYPE: 253bp Hindlll/EcoRV fragment + 343bp Hind
                                  11/EcoRl fragment
        (vi) ORIGINAL SOURCE:
                      (A) ORGANISM: whole cell
                      (B) STRAIN: CM4-184
                      (C) INDIVIDUAL ISOLATE: Z.Dai, et al
(D) DEVELOPMENTAL STAGE:
                      (E) HAPLOTYPE:
                      (F) TISSUE TYPE:
                      (G) CELL TYPE:
                      (H) CELL LINE:
                      (I) ORGANELLE:
        (vii) IMMEDIATE SOURCE:
                     (A) LIBRARY: genomic library of CM4-184
(B) CLONE: POS-1
        (viii) POSITION IN GENOME:
                      (A) CHROMOSOME/SEGMENT:
                      (B) MAP POSITION:
                      (C) UNITS:
        (ix)
               FEATURE:
                     (A) NAME/KEY: CaMV 35S promoter with duplication
                                       of upstream B domain
                     (B) LOCATION:(C) IDENTIFICATION METHOD:
                      (D) OTHER INFORMATION:
        (x)
               PUBLICATION INFORMATION:
                     (A) AUTHORS: (B) TITLE:
                      (C) JOURNAL:
                      (D) VOLUME:
                     (E) ISSUE: (F) PAGES:
                      (G) DATE:
                     (H) DOCUMENT NUMBER:
                     (I) FILING DATE:
                     (J) PUBLICATION DATE:
                      (K) RELEVANT RESIDUES IN SEQ ID NO:
        FROM (position) TO (position) (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 8:
CCC ACAGATGGTT AGAGAGGCTT ACGCAGCAGG TCTCATCAAG ACGATCTACC
CGAGCAATAA TCTCCAGGAA ATCAAATACC TTCCCAAGAA GGTTAAAGAT GCAGTCAAAA 113
GATTCAGGAC TAACTGCATC AAGAACACAG AGAAAGATAT ATTTCTCAAG ATCAGAAGTA 173
CTATTCCAGT ATGGACGATT CAAGGCTTGC TTCACAAACC AAGGCAAGTA ATAGAGATTG 233
GAGTCTCTAA AAAGGTAGTT CCCACTGAAT CAAAGGCCAT GGAGTCAAAG ATTCAAATAG 293
AGGACCTAAC AGAACTCGCC GTAAAGACTG GCGAACAGTT CATACAGAGT CTCTTACGAC 353
TCAATGACAA GAAGAAAATC TTCGTCAACA TGGTGGAGCA CGACACACTT GTCTACTCCA 413
AAAATATCAA AGATACAGTC TCAGAAGACC AAAGGGCAAT TGAGACTTTT CAACAAAGGG 473
TAATATCCGG AAACCTCCTC GGATTCCATT GCCCAGCTAT CTGTCACTTT ATTGTGAAGA 533
TAGTGGAAAA GGAAGGTGGC TCCTACAAAT GCCATCATTG CGATAAAGGA AAGGCCATCG 593
TTGAAGATGC CTCTGCCGAC AGTGGTCCCA AAGATGGACC CCCACCCACG AGGAGCATCG 653
TGGAAAAAGA AGACGTTCCA ACCACGTCTT CAAAGCAAGT GGATTGATGT GATAACATGG 713
TGGAGCACGA CACACTTGTC TACTCCAAAA ATATCAAAGA TACAGTCTCA GAAGACCAAA 773
GGGCAATTGA GACTTTTCAA CAAAGGGTAA TATCCGGAAA CCTCCTCGGA TTCCATTGCC 833
CAGCTATCTG TCACTTTATT GTGAAGATAG TGGAAAAGGA AGGTGGCTCC TACAAATGCC 893
ATCATTGCGA TAAAGGAAAG GCCATCGTTG AAGATGCCTC TGCCGACAGT GGTCCCAAAG 953
```

WO 98/21348 PCT/US97/20603

-29-

ATGGACCCCC	ACCCACGAGG	AGCATCGTGG	AAAAAGAAGA	CGTTCCAACC	ACGTCTTCAA	1013
AGCAAGTGGA	TTGATGTGAT	ATCTCCACTG	ACGTAAGGGA	TGACGCACAA	TCCCACTATC	1073
CTTCGCAAGA	CCCTTCCTCT	ATATAAGGAA	GTTCATTTCA	TTTGGAGAGA	ACACGGGGGA	1133
CTCTAGAGGA	TCCCCGGGTG	GTCAGT				1159

CLAIMS

We claim:

- 1. A method of producing human growth factors from plant cells, comprising the steps of:
 - (a) obtaining a positive transformant of the plant cells, the positive transformant carrying genetic material encoding the production of a human growth factor with a length of at least 200 amino acids;
 - (b) cultivating the positive transformant; and
- 10 (c) obtaining the human growth factors.
 - 2. The method as recited in claim 1, wherein obtaining the positive transformant has the step of:
- modifying a chimeric cDNA encoding the human growth factor with a length of at least 200 amino acids, for subcloning into a plant expression vector.
 - 3. The method as recited in claim 2, further comprising the steps of:
 - (a) subcloning the chimeric cDNA into the plant expression vector and obtaining a subcloned plant expression vector;
- 20 (b) transferring the subcloned plant expression vector into a plurality of plant cells;
 - (c) selecting a plurality of positive transformants from the plurality of plant cells on an antibiotic selective media;
- (d) permitting growth of the portion of the plurality of plant cells in whole plants or suspensions; and
 - (e) extracting a liquid containing the human growth factor from the plurality of transgenic plant cells.
- 4. The method as recited in claim 3, wherein transferring is by direct particle bombardment.

- 5. The method as recited in claim 3, wherein transferring is by Agrobacterium mediated transformation.
- 6. The method as recited in claim 5, wherein Agrobacterium mediated transformation comprises the steps of:
 - (a) placing the subcloned plant expression vector to an agrobacterium;
 - (b) co-cultivating the Agrobacterium containing the subcloned plant expression vector with the plurality of plant cells.
 - 7. The method as recited in claim 1, wherein the step of cultivating is with a whole plant.
- 8. The method as recited in claim 1, wherein the step of cultivating is with a plant tissue culture.
 - 9. The method as recited in claim 1, wherein the step of obtaining is selected from the group consisting of ultrafiltration, affinity chromatography, and electrophoresis.

The method as recited in claim 1, wherein the length of at least 200

amino acids is obtained by cloning a cDNA.

- 11. The method as recited in claim 10, wherein said cDNA is a pre-pro-25 EGF cDNA.
 - 12. The method as recited in claim 1, wherein the length of at least 200 amino acids is obtained by synthesizing a cDNA.

- 13. The method as recited in claim 12, wherein said synthesizing is concatomerizing multiple gene copies to obtain the length of at least 200 amino acids.
- 5 14. The method as recited in claim 1, further comprising increasing an overall size of a gene to be expressed with a fusion construct encoding an hEGF linked to a protein that is efficiently produced in plant systems.
- 15. The method as recited in claim 1, wherein said human growth factor is selected from the group consisting of epidermal growth factor (EGF), transforming growth factor (TGF), vascular endothelial growth factor (VEGF), platelet-derived growth factor (PDGF), fibroblast growth factor (FGF), tumor necrosis factor (TNF), heparin-binding epidermal growth factor (HBEGF), insulin-like growth factor (ILGF), platelet-derived endothelial cell growth factor (PDECGF), platelet-derived angiogenesis factor (PDAF), and bone-and-cartilage inducing growth factor (BCIF) and combinations thereof.
 - 16. The method as recited in claim 2, wherein modifying is by adding a regulatory element selected from the group consisting of leader sequences, signal peptides, transcription promoters or enhancers, and transcription terminators.
 - 17. The method as recited in claim 2, wherein modifying a chimeric cDNA, comprises the steps of:
- (a) adding said transcription promoter to the upstream or 5' end of the chimeric cDNA; and
 - (b) adding said transcription terminator to the downstream or 3' end of the chimeric cDNA.
- 18. The method as recited in claim 17, further comprising adding an additional regulatory element encoding a signal peptide, said additional regulatory

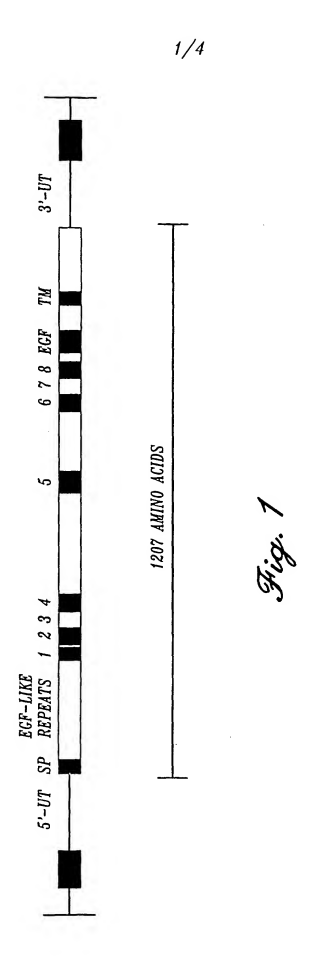
element added between the transcription promoter and the upstream 5' end of the chimeric cDNA.

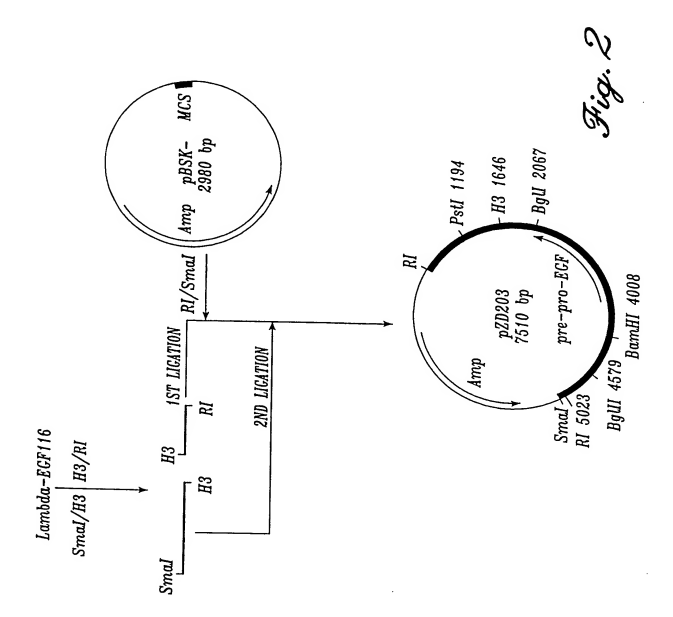
- The method as recited in claim 18, further comprising adding a
 regulatory element between the transcription promoter and the additional regulatory element encoding the signal peptide to enhance mRNA stability.
 - 20. The method as recited in claim 18, further comprising adding a regulatory element at the downstream or 3' end of the chimeric cDNA to enhance mRNA stability.
 - 21. The method as recited in claim 17, wherein transcription promoters limit growth factors production to a non-crop portion of a transgenic whole plant.
- 15 22. The method as recited in claim 21, wherein the transcription promoters are selected from the group consisting of an upstream enhancer region (-343 to -90 bp) of a CaMV 35S promoter, a chlorophyll a/b binding promoter (cab1) and combinations thereof.
- 20 23. The method as recited in claim 17, wherein the transcription promoters are selected from the group consisting of a modified 35S promoter, TSC29 promoter, TSC40 promoter and combinations thereof.
- 24. The method as recited in claim 23, wherein the modified 35S

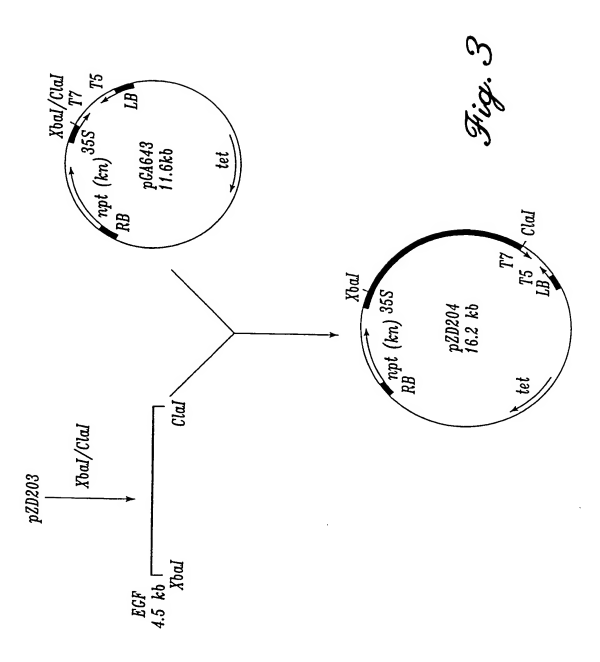
 25 promoter is a 35S promoter modified by duplicating an upstream enhancer region

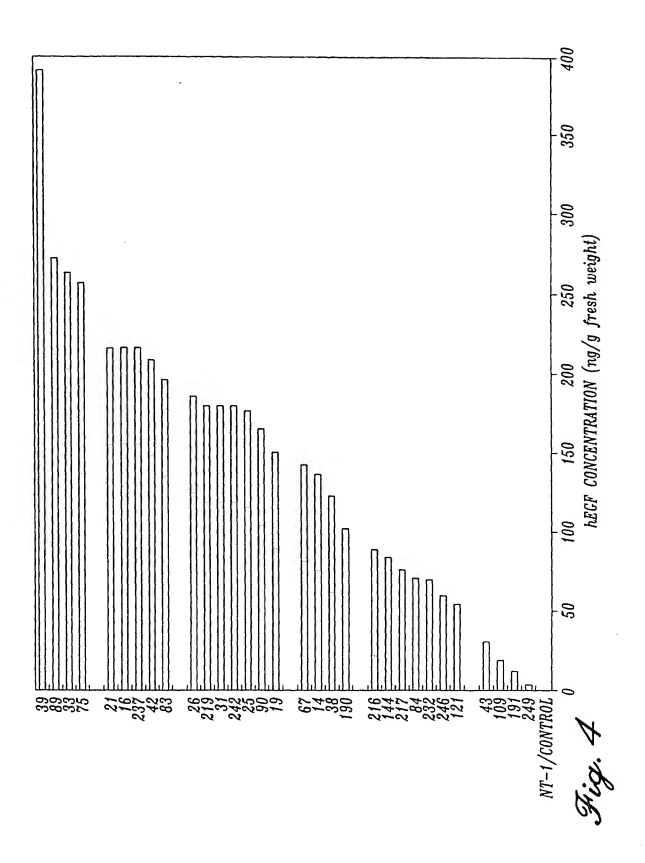
 (-343 to -90 bp) of the 35S promoter to increase transcription activity.
 - 25. The method as recited in claim 2, wherein said cDNA is a pre-pro-EGF cDNA.

- 26. The method as recited in claim 25, wherein said pre-pro-EGF cDNA has approximately 4.5 kb, whereby overall titers of active hEGF in both whole plants and cell culture are increased.
- 5 27. The method as recited in claim 2, wherein the length of at least 200 amino acids is obtained by synthesizing the cDNA.
- 28. The method as recited in claim 27, wherein said synthesizing is concatomerizing multiple gene copies to obtain the length of at least 200 amino acids.
 - 29. The method as recited in claim 28, wherein said multiple gene copies are an oligomeric polypeptide having of repeated hEGF domains.
- 15 30. The method as recited in claim 2, further comprising increasing an overall size of a gene to be expressed with a fusion construct encoding an hEGF linked to a protein that is efficiently produced in plant systems.









INTERNATIONAL SEARCH REPORT

information on patent family members

Inter anal Application No
PCT/US 97/20603

			3 97/20003
Patent document cited in search report	Publication date	Patent family member(s)	Publication date
WO 8700865 A	12-02-87	US 4956282 A AU 603063 B AU 6196686 A DE 3687705 A EP 0233915 A JP 63500425 T US 5550038 A US 5629175 A	11-09-90 08-11-90 05-03-87 18-03-93 02-09-87 18-02-88 27-08-96 13-05-97
WO 8903887 A	05-05-89	CA 1337048 A AU 2811889 A AU 634987 B AU 4495189 A CA 2000661 A WO 9004032 A EP 0319353 A EP 0318341 A JP 2501802 T JP 3502644 T US 5487991 A AT 143694 T AT 140959 T DE 3855455 D DE 3855591 D DE 3855591 T EP 0723019 A US 5623067 A	19-09-95 23-05-89 11-03-93 01-05-90 14-04-90 07-06-89 31-05-89 21-06-90 20-06-91 30-01-96 15-10-96 15-08-96 05-09-96 09-01-97 07-11-96 03-04-97 24-07-96 22-04-97
WO 9514099 A	26-05-95	US 5693506 A AU 1289295 A CA 2176834 A EP 0788550 A JP 9509565 T	02-12-97 06-06-95 26-05-95 13-08-97 30-09-97
WO 9321320 A	28-10-93	AU 1583292 A AU 678154 B AU 3884493 A EP 0636179 A EP 0633940 A	18-11-93 22-05-97 08-11-93 01-02-95 18-01-95

INTERNATIONAL SEARCH REPORT

Information on patent family members

Inte .onal Application No
PCT/US 97/20603

Patent document cited in search report	Publication date	ı	Patent family member(s)	Publication date
WO 9321320 A		FI FI JP NO NO WO US	944550 A 944841 A 7507199 T 7505525 T 943664 A 943914 A 9320216 A 5650554 A	30-11-94 14-12-94 10-08-95 22-06-95 15-11-94 09-12-94 14-10-93 22-07-97

Form PCT/ISA/210 (patent family annex) (July 1992)





